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(54)	COMPLEXED LIQUID FUEL COMPOSITIONS				
(75)	Inventor:	John P. Consaga, La Plata, MD (US)			
(73)	Assignee:	The United States of America as represented by the Secretary of the Navy, Washington, DC (US)			
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(52)	U.S. Cl	C06D 5/00 			
(58)	Field of Se	earch 149/88, 108; 60/205, 60/217			

### (56)References Cited

### U.S. PATENT DOCUMENTS

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			Camp et al
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5,114,506			Consaga et al 149/88
5,398,612	*	3/1995	Graham et al 102/287
5,440,993		8/1995	Osofsky 102/374
5,639,987		6/1997	Berteleau et al 149/19.8
5,652,409		7/1997	Thompson et al 149/98

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Primary Examiner—Michael J. Carone Assistant Examiner—Glenda L Sánchez (74) Attorney, Agent, or Firm-Mark Homer

### (57)ABSTRACT

A complexed liquid fuel composition has the components of a cyclodextrin nitrate, a nitrate ester plasticizer, bismuth subsalicylate and a stabilizer.

18 Claims, No Drawings

## COMPLEXED LIQUID FUEL COMPOSITIONS

This application is a division of application Ser. No. 09/443,120, filed Nov. 18, 1999 now U.S. Pat. No. 6,110,

## SPONSORED RESEARCH OR DEVELOPMENT STATEMENT REGARDING FEDERALLY

any royalties thereon or therefor. used by or for the government of the United States of America for governmental purposes without the payment of The invention described herein may be manufactured and

# BACKGROUND OF THE INVENTION

## Field of the Invention

as a propellant. an appropriate energy, stability and sensitivity that is useful subsalicylate and stabilizer to form liquid compositions with particularly, the organic nitrate esters of the present inven-tion are complexed with a nitrate ester plasticizer, bismuth contain an energetic material of organic nitrate esters. Most particularly, the liquid propellants of the present invention The present invention relates to liquid propellants. More

# Brief Description of the Related Art

No. 5,652,409 to Thompson et al. discloses a non-complexed solid double-based propellant having cyclodex-trin nitrate, nitroglycerin, and bismuth salicylate. However, the identified explosive compositions are not complexed, nents to be suitable as a liquid fuel composition. Berteleau et al. discloses a solid propellant containing cellulose nitrate, nitroglycerin and bismuth salicylate. U.S. Pat. No. 5,652,409 to Thompson et al. discloses a nongun propellant or explosive composite having a solid nitrate ester of cyclodextrin and nitroglycerin. U.S. Pat. No. 5,440, 993 to Osofsky discloses a high velocity rocket containing nitroglycerin and nitrocellulose. U.S. Pat. No. 5,639,987 to lack sufficiently stability and/or lack the requisite compo-Pat. No. 5,114,506 to Consaga et al. discloses an energetic Several types of energetic compositions are known. U.S.

present invention addresses this need In view of the foregoing, there is a need for a highly energetic material useful as a liquid fuel propellant. The

# SUMMARY OF THE INVENTION

energetic compound. the cyclodextrin nitrate, nitrate ester plasticizer, bismuth subsalicylate and stabilizer are complexed together into an plasticizer, bismuth subsalicylate and a stabilizer, wherein composition comprising a cyclodextrin nitrate, a nitrate ester The present invention includes a complexed liquid fuel 80

sion comprising the steps of inputing into a combuster a composition in the combuster, wherein thrust is produced and a stabilizer, and reacting the complexed liquid fuel trin nitrate, a nitrate ester plasticizer, bismuth subsalicylate complexed liquid fuel composition comprising a cyclodex-The present invention also includes a method for propul-

## DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENT

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ers. The composition is used in a method for propulsion that complexed compositions containing cyclodextrin nitrates, nitrate ester plasticizers, bismuth subsalicylate and stabilizrials useful as liquid propellants. The present invention relates generally to energetic mate-als useful as liquid propellants. The energetic materials are

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allows an appropriate energy, stability and sensitivity that is useful as a liquid propellant.

S separate. The free movement liquid characteristics of the present invention provide flow, and may be pumped, from one location into another, such as into a combuster. Liquid fuel compositions of the present invention provide "free movement" of the composition without a tendency to is slightly higher than water. Generally, the viscosity of the liquid of the present invention

4 35 30 25 20 15 10 period of from about 1 to about 4 hours. energy is preferably in the form of shear mixing, using shear blades to mix the composition. The acetone medium permits the components to dissolve, particularly the cyclodextrin 140° F. or higher, and pressures of from about 25–30 mm Hg that are continuously decreased to about 3 mm Hg over a Preferably, acetone is used with temperatures of from about nitrates. As low elevated temperatures strip the medium cal energy is placed into the component parts. Mechanical elevated temperature, with the medium selected for its ability to dissolve the components and be removed at modest characteristics of each component. Complexing may be imparted into the composition of the present invention with the addition of heat and mechanical energy, i.e., shear, under component parts of the composition, i.e., the cyclodextrin nitrates, nitrate ester plasticizers, bismuth subsalicylate and stabilizer are "tied" to one another within the complexed composition. As such, the component parts of the compomuth subsalicylate and stabilizer are processed into complexed energetic compositions of the present invention. Complexed compositions include an intermolecular the shearing complexes the components in the composition. from the mixed components in an evacuated environment, complexing components. Vacuum is applied while mechanitemperatures, i.e., temperatures that are not damaging to the are mixed together in acetone or other like medium at an vacuum, in an appropriate medium, such as acetone. For example, the individual components of the present invention not complexed within a composition retain the individual raised boiling point. By contrast, mixed components that are may be evidenced by composition characteristics, such as a sition tend to act as a single ingredient or material, which attraction, i.e., dipole-The cyclodextrin nitrates, nitrate ester plasticizers, bis the component parts of the compo--dipole or ion-dipole, between the

55 \$ Different  $\alpha$ -cyclodextrin mitrate esters, based on the same basic  $\alpha$ -cyclodextrin moiety, differ from each other in the Preferably an average of from about 2 to about 3, more preferably from about 2.5 to about 3, and most preferably from about 2.6 to about 3 nitrate ester groups (—ONO<sub>2</sub>) per D-glucose unit are present in the mitration product of the degree of nitration, i.e., nitrate ester unit content. Likewise, ester, either individually or within various mixtures thereof α-cyclodextrin, β-cyclodextrin or γ-cyclodextrin nitrate capable of being nitrated to a nitrate ester group of has 21—OH groups, and significantly greater increase over α-cyclodextrin with 18—OH groups. Each D-glucose unit sesses a larger cavity, allowing for approximately an 80% merease in cavity size from β-cyclodextrin nitrate, which nitrate. γ-cyclodextrin nitrate is particularly desirable because the maximum energy potential of the γ-cyclodextrin nitrate is significantly higher than other cyclodextrin nitrate in a cyclodextrin compound has three free γ-cyclodextrin nitrate, with 24 available compounds, while it retains significant stability. include energetic materials such as a-cyclodextrin, preferred cyclodextrin nitrate comprises γ-cyclodextrin  $\beta$ -cyclodextrin,  $\gamma$ -cyclodextrin, and mixtures thereof. The Cyclodextrin nitrate compounds of the present invention -OH groups, pos-OH groups  $ONO_{2}$ 

different  $\beta$ -cyclodextrin nitrate esters differ from each other in the degree of nitration, as do different  $\gamma$ -cyclodextrin nitrate esters.

The cyclodextrins of the present invention may be nitrated using conventional techniques that are used in the preparation of nitrocellulose, with the degree of nitration controlled by varying the nitration conditions. Formation of the cyclodextrins is disclosed in U.S. Pat. No. 5,114,506 to Consaga et al., issued May 19, 1992, the disclosure of which is herein incorporated by reference. Commercial γ-cyclodextrins are available from Wacker-Bio-chem of Edieville, Iowa under the tradename Cavamax-W8.

The cyclodextrin nitrate esters of the present invention provide useful replacements for energetic organic nitrate ester plasticizers within the liquid fuel composition as the cyclodextrin nitrate esters increase the thermal stability and decrease the shock sensitivity of the liquid fuel composition of the organic nitrate ester plasticizers. The cyclodextrin nitrate esters also possess comparable or greater energy content than the organic nitrate esters are sensitive to electrostatic discharge (ESD), e.g., β-cyclodextrin nitrate ester granticizers. As dry powders, the cyclodextrin nitrate esters are sensitive to electrostatic discharge (ESD), e.g., β-cyclodextrin nitrate ester (β-CDN) (C<sub>42</sub>H<sub>32</sub>N<sub>18</sub>O<sub>71</sub>) has an ESD value of only 0.0125 joules. When the organic nitrate ester plasticizer of 1.1.1-trimethylolethane trinitrate (TMETN), having an ESD value of 12.5 joules, is mixed with β-CDN (2:1 weight ratio), the resulting composite mixture has a liquid consistency and a resultant ESD value of 12.5 joules. The composite mixture, however, has a low shock sensitivity.

The cyclodextrin starting materials comprise cyclic structures having  $1,4-\alpha$ -glucosidically linked D-glucose units, preferably being  $\alpha$ -cyclodextrin with  $6,\beta$ -cyclodextrin with  $7,\gamma$ -cyclodextrin with 8 glucosidically linked D-glucose units, or mixtures of these compounds. A preferred embodiment of the present invention comprises an energetic composite comprising a nitrate ester of  $\gamma$ -cyclodextrin with a majority of the —OH groups fully nitrated, and an organic nitrate ester plasticizer of 1,1,1-trimethylolethane trinitate. Preferably, the weight ratio of the 1,1,1-trimethylolethane trinitate to the nitrate ester of  $\gamma$ -cyclodextrin ranges from about 2:1 to about 6:1 or less, and more preferably from about 2:1 to about 5:1.

The cyclodextrin nitrate preferably comprises from about 20 wt % to about 50 wt % of the complexed liquid fuel composition, more preferably from about 25 wt % to about 40 wt %, and most preferably approximately 32.5 wt % of the complexed liquid fuel composition.

Suitable nitrate ester plasticizers of the present invention that are complexed with the cyclodextrin nitrate are determinable by those skilled in the art, by considering the energy potential and sensitivity desired. Preferred energetic organic nitrate ester plasticizers include 1,1,1-trimethylolethane trinitate (TMETN), 1,2,4-butanetriol trinitrate (BTIN), triethylene glycol dinitrate (TEGDN), nitroglycerin (NG), 1,2-propyleneglycol dinitrate (PGDN), pentacrythritol trinitrate (PETRIN), diethylene glycol dinitrate (PGDN), and combination or mixtures of these compounds. More preferred energetic organic nitrate ester plasticizers include the individual compounds or mixtures of 1,1,1-trimethylolethane trinitrate, 1,2,4-butanetriol trinitrate, triethylene glycol dinitrate, and nitroglycerin. Nitroglycerin is most preferred, which is commercially available from Naval Surface Warfare Center, Indian Head, Md.

Operable amounts of cyclodextrin nitrate ester to ener- 65 getic organic nitrate ester plasticizer vary with the choice of cyclodextrin nitrate ester and energetic nitrate ester

plasticizer, but generally range from about 1:1 to about 1:6 with amounts of 1:2, 1:3 and 1:4 operable with at least enough plasticizer to convert the powdery cyclodextrin nitrate ester into a liquid composition. With the combination of the cyclodextrin nitrate ester and nitrate ester plasticizer, the ESD of the nitrate ester plasticizer decreases to about that of the cyclodextrin nitrate ester while retaining the low shock sensitivity of the cyclodextrin nitrate ester plasticizer cause a saturation point to be reached, after which the plasticizer remains separate or neat, i.e., not complexed, from the composition with the neat plasticizer retaining high shock sensitivity.

Generally, the amount of nitrate ester plasticizer ranges is from about 50 wt % to about 80 wt % of the complexed liquid fuel composition, with amounts of from about 60 wt % to about 75 wt % more preferred, and an amount of approximately 65 wt % of the complexed liquid fuel composition most preferred.

Bismuth subsalicylate is an acetone soluble complexing component with the cyclodextrin nitrate ester and nitrate ester plasticizer that provides a burn rate modifier to the liquid fuel composition and a complex stabilizer. As such, the bismuth subsalicylate inhibits the breakup of the liquid fuel composition into its component parts. This imparts significant safety to the liquid fuel composition in storage, handling and manufacturing. Preferably, the bismuth subsalicylate comprises from about 0.75 wt % to about 1.5 wt % of the complexed liquid fuel composition. Bismuth subsalicylate is commercially available from Pfaltz & Bauer, Inc. of Waterburg, Conn.

The stabilizer component of the present invention comprises a stabilizing compound having a pH of from about 7 or less to ensure decomposition of the nitrate ester does not occur. Preferably, the stabilizer comprises an acidic or neutral amide, with more preferred stabilizers including 2-nitrodiphenyl amine (2NDPA), methylnitroaniline (MNA) and/or combinations thereof. Preferred amounts of stabilizer range from about 1 wt % to about 2 wt % of the complexed liquid fuel composition.

Increases in the amount of cyclodextrin nitrate ester, bismuth subsalicylate and/or stabilizer in relation to the nitrate ester plasticizer on average cause a decrease in the amount of available energy of the liquid fuel composition. The appropriate relative amounts of these components for a particular liquid fuel composition is determinable by those skilled in the art, generally as a factor of the liquidity and available energy of the complexed composition. As additional components tend to decrease the available energy within the complexed components, other energetic and nonenergetic components generally are not added to control the liquidity and available energy of the complexed composition.

55 Propulsion is created by inputting the complexed liquid finel composition into a combuster, reacting the complexed liquid finel composition, and allowing a controlled release of the reaction product. Combusters are known in the art, generally comprising any suitable reaction chamber for designed for propulsion with the reaction of a highly energetic chemical composition, including reaction chambers for missiles, rockets, space vehicles, and other such propelling apparatuses. Reacting may be accomplished by an ignition or heat source sufficient for the complexed composition to initiate continuous combustion. Combustion is maintained with the continuous feeding of unreacted complexed liquid fuel composition into the combuster. Proper release of the

resulting gases from the combustion within the combuster provides thrust.

The complex of the cyclodextrin nitrate, nitrate ester plasticizer, bismuth subsalicylate and stabilizer provides a particularly suitable liquid fuel for a limited fuel source for extended flight. Combinations of the γ-cyclodextrin nitrate and nitroglycerin complexed with the bismuth subsalicylate and stabilizer are particularly useful for a large energy source from a relatively small amount of composition. Compositions range from about 25 wt % to about 40 wt % γ-cyclodextrin nitrate, from about 60 wt % to about 75 wt % nitroglycerin, from about 1 wt % to about 2 wt % bismuth subsalicylate and from about 1 wt % to about 2 wt % stabilizer. Most preferred liquid fuel compositions include approximately 32.5 wt % γ-cyclodextrin nitrate, 65 wt % nitroglycerin, 1.05 wt % bismuth subsalicylate and 1.4 wt % nitroglycerin, 1.05 wt % bismuth subsalicylate and 1.4 wt % NIDPA

The following examples illustrate suitable combinations of the cyclodextrin nitrate, nitrate ester plasticizer, bismuth subsalicylate and stabilizer of the present invention that are expected to provide highly energetic liquid fuel compositions.

### **EXAMPLE** 1

with the pressure gradually decreased to 3 mm Hg by the end of the first 60 minutes, with the pressure maintained at 3 mm Hg through the second 60 minute period. The resulting liquid is analyzed under thermogravimetric analysis (TGA) wt % 2-nitrodiphenyl amine), and 11 grams of bismuth subsalicylate are added with 3 liters of acetone, and heated and stirred for 30 minutes at a temperature of 140° F. under a pressure of 20 mm Hg. The second step is repeated a third Baker-Perkins vertical mixer, manufactured by Baker-Perkins of Saginaw, Mich., for 30 minutes at a temperature of 140° F. under a pressure of 20 mm Hg. In a second step, and fourth time under similar conditions. After the fourth step, the mixture is maintained at 140° F. for 120 minutes the actione, with the nitroglycerin that normally peaks at 130-135°F, expected to peak at approximately 165-170°F. 2-nitrodiphenyl amine), 670 grams of nitroglycerin (with 1 another 340 grams of y-cyclodextrin (with 1 wt which is included in the acetone. The mixture is stirred in a salicylate in 3 liters of acetone. The γ-cyclodextrin and In a first step, 340 grams of γ-cyclodextrin is mixed with 670 grams of nitroglycerm, and 11 grams of bismuth subto verify complexation of the composition and removal of nitroglycerin are stored in 1 wt % 2-nitrodiphenyl amine 2-nitrodiphenyl ક 6 35

### **EXAMPLE 2**

In a first step, 350 grams of y-cyclodextrin is mixed with 630 grams of 1,1,1-trimethylolethane trinitrate, and 10 grams of bismuth subsalicylate in 2.5 liters of acetone. The y-cyclodextrin and 1,1,1-trimethylolethane trinitrate are stored in 1 wt % 2-nitrodiphenyl amine which is included in 55 the acetone. The mixture is stirred for 30 minutes at a temperature of 150° F. under a pressure of 20 mm Hg. In a second step, another 350 grams of y-cyclodextrin (with 1 wt % 2-nitrodiphenyl amine), 630 grams of 1,1,1-trimethylolethane trinitrate (with 1 wt % 2-nitrodiphenyl amine), and 10 grams of bismuth subsalicylate are added with 2.5 liters of acetone, and heated and stirred for 30 minutes at a temperature of 150° F. under a pressure of 20 mm Hg. The second step is repeated a third and fourth time under similar conditions. After the fourth step, the mixture 65 maintained at 150° F. for 120 minutes with the pressure gradually decreased to 3 mm Hg by the end of the first 60

minutes, with the pressure maintained at 3 mm Hg through the second 60 minute period. The resulting liquid is analyzed under thermogravimetric analysis (TGA) to verify complexation of the composition and removal of the acetone, with the 1,1,1-trimethylolethane trinitrate that normally peaks at 133° F, expected to peak at approximately 165–170° F.

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### **EXAMPLE 3**

30 20 15 10 to verify complexation of the composition and removal of the acetone, with the 1,2,4-butanetriol trinitrate that normally peaks at 135–140° F, expected to peak at approxi-140° F. under a pressure of 20 mm Hg. In a second step, another 340 grams of  $\alpha$ -cyclodextrin (with 1 wt % methylnitroaniline), 670 grams of 1,2,4-butanetriol trinitrate (with 1 wt % methylnitroaniline), and 11 grams of bismuth and stirred for 30 minutes at a temperature of 140° F. under a pressure of 20 mm Hg. The second step is repeated a third mately 165-170° F. liquid is analyzed under thermogravimetric analysis (TGA) Hg through the second 60 minute period. The resulting of the first 60 minutes, with the pressure maintained at 3 mm with the pressure gradually decreased to 3 mm Hg by the end and fourth time under similar conditions. After the fourth step, the mixture is maintained at 140° F. for 120 minutes subsalicylate are added with 3 liters of acetone, and heated wt % methylnitroaniline which is included in the acetone. α-cyclodextrin and 1,2,4-butanetriol trinitrate are stored in 1 670 grams of 1,2,4-butanetriol trinitrate, and 11 grams of bismuth subsalicylate in 3 liters of acctone. The The mixture is stirred for 30 minutes at a temperature of In a first step, 340 grams of \alpha-cyclodextrin is mixed with

### **EXAMPLE 4**

In a first step, 340 grams of β-cyclodextrin is mixed with 670 grams of TMETN, and 11 grams of bismuth subsalicy-late in 3 liters of acetone. The β-cyclodextrin and triethylene glycol dinitrate are stored in 1 wt % methylnitroaniline which is included in the acetone. The mixture is stirred for 30 minutes at a temperature of 140° F. under a pressure of β-cyclodextrin (with 1 wt % methylnitroaniline), 670 grams of β-cyclodextrin (with 1 wt % methylnitroaniline), and 11 grams of bismuth subsalicylate are added with 3 liters of acetone, and heated and stirred for 30 minutes at a temperature of 140° F. under a pressure of 20 mm Hg. The second step is repeated a third and fourth time under similar conditions. After the fourth step, the mixture is maintained at 140° F. for 120 minutes with the pressure gradually decreased to 3 mm Hg by the end of the first 60 minutes, with the pressure maintained at 3 mm Hg through the second 60 minute period. The resulting liquid is analyzed under thermogravimetric analysis (TGA) to verify complexation of the composition and removal of the acetone, with the TMETN that normally peaks at 130–135° F., expected to peak at approximately 165–170° F.

The foregoing summary, description, and examples of the invention are not intended to be limiting, but are only exemplary of the inventive features which are defined in the claims.

What is claimed is:

 A method of propulsion comprising the steps of: inputing into a combuster a complexed liquid fuel composition comprising a cyclodextrin nitrate, a nitrate ester plasticizer, a bismuth subsalicylate and a stabi-

reacting the complexed liquid fuel composition in the combuster, wherein thrust is produced.

lizer; and,

- The method of claim 1, wherein the cyclodextrin nitrate comprises γ-cyclodextrin nitrate.
   The method of claim 1, wherein the nitrate ester
- plasticizer comprises nitroglycerin.
- 4. The method of claim 1, wherein the complexed liquid fuel comprises from about 25 wt % to about 40 wt % y-cyclodextrin nitrate, and from about 60 wt % to about 75 % nitroglycerin
- 5. The method of claim 4, wherein the complexed liquid fuel comprises approximately 32.5 wt % \(\gamma\)-cyclodextrin nitrate and approximately 66 wt % nitroglycerin.
  6. The method of claim 1, wherein the complexed liquid fuel comprises approximately 32.5 wt % \(\gamma\)-cyclodextrin nitrate, 66 wt % nitroglycerin and 1.05 wt % bismuth 10
- subsalicylate

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- comprises an energetic material selected from the group consisting of  $\alpha$ -cyclodextrin,  $\beta$ -cyclodextrin,  $\gamma$ -cyclodextrin, and mixtures thereof. 7. The method of claim 1, wherein the cyclodextrin nitrate
- comprises from about 20 wt % to about 50 wt % of the 8. The method of claim 1, wherein the cyclodextrin nitrate 20
- complexed liquid fuel composition.

  9. The method of claim 8, wherein the cyclodextrin nitrate comprises from about 25 wt % to about 40 wt % of the complexed liquid fuel composition.
- nitrate comprises approximately 32.5% of the complexed 10. The method of claim 9, wherein the cyclodextrin
- liquid fuel composition.

  11. The method of claim 1, wherein the nitrate ester plasticizer comprises an energetic material selected from the group consisting of 1,1,1-trimethylolethane trinitate (TMEIN), 1,2,4-butanetriol trinitrate (BTIN), triethylene (TMEIN), 1,2,4-butanetriol trinitrate (BTIN), triethylene glycol dinitrate (TEGDN), nitroglycerin (NG), 1,2-30

- propyleneglycol dinitrate (PGDN), pentaerythritol trinitrate (PETRIN), diethylene glycol dinitrate (DEGN), and mix-
- tures thereof.

  12. The method of claim 1, wherein the nitrate ester plasticizer comprises from about 50 wt % to about 80 wt % of the complexed liquid fuel composition.

  13. The method of claim 1, wherein the nitrate ester

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- plasticizer comprises from about 60 wt % to about 75 wt %
- of the complexed liquid fuel composition.

  14. The method of claim 1, wherein the nitrate ester plasticizer comprises approximately 65 wt % of the complexed liquid fuel composition.
- 15. The method of claim 1, wherein the bismuth subsali-

cylate comprises from about 0.75 wt % to about 1.5 wt % of

- the complexed liquid fuel composition.

  16. The method of claim I, wherein the stabilizer comprises a stabilizing compound having a pH of from about 7 or less selected from the group consisting of 2-nitrodiphenyl thereof amine (2NDPA), methylnitroaniline (MNA), and mixtures
- 17. The method of claim 1, wherein the stabilizer comprises from about 1 wt % to about 2 wt % of the complexed liquid fuel composition.
- comprising the steps of: 18. Thrust to propel a rocket produced from the process

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- inputing into a combuster a complexed liquid fuel comlizer; and, position comprising a cyclodextrin nitrate, a nitrate ester plasticizer, a bismuth subsalicylate and a stabi-
- reacting the complexed liquid fuel composition in the combuster